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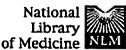
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	DB=PC	GPB, USPT, USOC, EPAB, JPAB, DWPI, TDBD; PLUR = YES; OP = OR	
	L34	L33 and APO-3	34
	L33	L32 and apoptosis	172
	L32	130 and antagonist	270
	L31	L30 and antaognist	0
	L30	(APO)same(antibod?)	952
	L29	L28 and apo-3	2.
	L28	424/178.1.ccls.	1051
	L27	L26 and APO-3	0
	L26	424/142.1.ccls.	241
	L25	L24 and Apo-3	1
	L24	424/133.1.ccls.	644
	L23	L22 and APO-3	3
	L22	424/130.1.ccls.	1634
	L21	L18 and APO-3	0
	L20	L18 and Apo-3	0
	L19	L18 and Apo-3 antibody	217182
	L18	435/7.21.ccls.	2559
	L17	435/7.21,ccls.	0
口	L16	L15 and Apo-3 receptor	209375
	L15	L14 and activation	47
	L14	L13 and apoptosis	47
	L13	L12 and inhibit	47
	L12	L11 and antibod?	47
	L11	L10 and Apo-3	47
	L10	435/7.1.ccls.	10005
	L9	L8 and anti-Apo-3	9
	L8	L7 and APO-3	157
	L7	(ashkenazi)	2569
	L6	(Avi)adj(J)adj(ashkenazi)	0
	L5	(A)adj(ashkenzai)	0

	L4	(ashkenzai)same(APO-3)	0
	DB=U	SPT; PLUR=YES; OP=OR	
	L3	US-6469144-B1.did.	1
•	DB=P	GPB, USPT, USOC, EPAB, JPAB, DWPI, TDBD; PLUR=YES; OP=OR	
	L2	(APO-3)same(antibod?)	19
	L1	(APO-3)same(antibod?)same(antagonist)same(apoptosis)same(NF)adj (kappa)adi(B)adi(activation)	0

END OF SEARCH HISTORY









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Related Articles, Links

Monoclonal antibody-mediated tumor regression by induction of apoptosis.

Trauth BC, Klas C, Peters AM, Matzku S, Moller P, Falk W, Debatin KM, Krammer PH.

Institute for Immunology and Genetics, German Cancer Research Center, Heidelberg.

To characterize cell surface molecules involved in control of growth of malignant lymphocytes, monoclonal antibodies were raised against the human B lymphoblast cell line SKW6.4. One monoclonal antibody, anti-APO-1, reacted with a 52-kilodalton antigen (APO-1) on a set of activated human lymphocytes, on malignant human lymphocyte lines, and on some patient-derived leukemic cells. Nanogram quantities of anti-APO-1 completely blocked proliferation of cells bearing APO-1 in vitro in a manner characteristic of a process called programmed cell death or apoptosis. Cell death was preceded by changes in cell morphology and fragmentation of DNA. This process was distinct from antibody- and complement-dependent cell lysis and was mediated by the antibody alone. A single intravenous injection of anti-APO-1 into nu/nu mice carrying a xenotransplant of a human B cell tumor induced regression of this tumor within a few days. Histological thin sections of the regressing tumor showed that anti-APO-1 was able to induce apoptosis in vivo. Thus, induction of apoptosis as a consequence of a signal mediated through cell surface molecules like APO-1 may be a useful therapeutic approach in treatment of malignancy.

PMID: 2787530 [PubMed - indexed for MEDLINE]

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☐ 1: Curr Biol. 1996 Dec 1;6(12):1669-76.

Related Articles, Links

Apo-3, a new member of the tumor necrosis factor receptor family, contains a death domain and activates apoptosis and NF-kappa B.

Marsters SA, Sheridan JP, Donahue CJ, Pitti RM, Gray CL, Goddard AD, Bauer KD, Ashkenazi A.

Department of Molecular Oncology, Genentech, Inc., South San Francisco, California 94080-4918, USA.

BACKGROUND: Two receptors that contain the so-called "death domain" have been described to date: tumor necrosis factor receptor 1 (TNFR1) and Fas/Apo-1 (CD95); both belong to the TNFR gene family. The death domain of TNFR1 mediates the activation of programmed cell death (apoptosis) and of the transcription factor NF-kappa B, whereas the death domain of CD95 only appears to activate apoptosis. RESULTS: We have identified an additional member of the TNFR family, which we have named Apo-3. Apo-3 is a transmembrane protein of approximately 47 kDa that has similarity of members of the TNFR family in its extracellular, cysteine-rich domains. In addition, Apo-3 resembles TNFR1 and CD95 in that it contains a cytoplasmic death domain. The Apo-3 gene mapped to human chromosome 1p36.3, and Apo-3 mRNA was detected in several human tissues, including spleen, thymus, peripheral blood lymphocytes, small intestine and colon. Ectopic expression of Apo-3 in HEK293 or HeLa cells induced marked apoptosis. CrmA, a poxvirus inhibitor of Ced-3-like proteases which blocks death signaling by TNFR1 and CD95, inhibited Apo-3-induced apoptosis. Ectopic expression of Apo-3 also induced the activation of NF-kappa B. Apo-3 did not specifically bind to the Apo-2 ligand, suggesting the existence of a distinct ligand for Apo-3. CONCLUSIONS: These results identify Apo-3 as a third member of the TNFR family that activates apoptosis, and suggest that Apo-3, TNFR1 and CD95 engage a common apoptotic cell-death machinery. Apo-3 resembles TNFR1 because it can stimulate NF-kappa B activity and regulate apoptosis. Apo-3 mRNA is expressed in various tissues, consistent with the possibility that this receptor may regulate multiple signaling functions.